application Serial No. 10/008,516, filed November 8, 2001, now US Patent No. 6,649,607."

As requested by the Examiner, Applicants have amended claim 32 so that claim 32 is a method of administration claim describing the dose at which a pharmaceutical composition comprising S-tofisopam is being administered.

## REJECTIONS

Claims 1-5, 28 and 29 have been rejected under 35 U.S.C. 102(a) as being anticipated by the Landry et al reference. The Landry reference teaches the use of (R)-tofisopam for the prevention and treatment of anxiety and anxiety disorders. (R) - tofisopam was found to be the active isomer of racemic tofisopam in the head twitch assay described in column 21, lines 24-34. (S)-tofisopam was used in the assay merely to show that the (S)-enantiomer was inactive in the assay. Thus, the Landry reference does not anticipate the present invention which describes (S)-tofisopam as an active pharmaceutical ingredient which can be used to effectively treat a disease.

The claims have been further amended to include the limitation that the composition that is being

envisioned is one that is pharmaceutically active, rather than a composition that is not pharmaceutically effective.

Furthermore, claim 29 has been amended to delete the language "approximately". None of these amendments introduces new matter.

## Double Patenting Rejections

The Examiner has provisionally rejected claims 1-5 and 28-32 under the judicially created doctrine of obviousness-type double patenting as being unpatentable over claim 31 of copending Application No. 10/781,422.

Applicants are submitting a terminal disclaimer to obviate the rejection.

The Examiner has nonprovisionally rejected claims

1-5 and 28-32 under the judicially created doctrine of
obviousness-type double patenting as being unpatentable

over claims 1-5 of US Patent No. 6,649,607. Applicants are
submitting a terminal disclaimer to obviate the rejection.

Applicants kindly request that the claims be accepted in view of the remarks and amendments provided above.

Respectfully submitted,

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## CLAIMS WITH MARKUPS

- 1. (Currently amended) A <u>pharmaceutical</u> composition comprising a therapeutically effective amount of Stofisopam, a prodrug or pharmaceutically acceptable salt thereof, substantially free of its R-enantiomer, with a pharmaceutically acceptable carrier.
- 2. (Currently amended) The <u>pharmaceutical</u> composition of claim 1 wherein the amount of S-tofisopam or a prodrug or a pharmaceutically acceptable salt thereof is 85% or more by weight of the total weight of tofisopam.
- 3. (Currently amended) The <u>pharmaceutical</u> composition of claim 1 wherein the amount of S-tofisopam or a prodrug or a pharmaceutically acceptable salt thereof is 90% or more by weight of the total weight of tofisopam.
- 4. (Currently amended) The <u>pharmaceutical</u> composition of claim 1 wherein the amount of S-tofisopam or a prodrug or a pharmaceutically acceptable salt thereof is 95% or more by weight of the total weight of tofisopam.

- 5. (Currently amended) The <u>pharmaceutical</u> composition of claim 1 wherein the amound of S-tofisopam or a prodrug or a pharmaceutically acceptable salt thereof is 99% or more by weight of the total weight of tofisopam.
- 28. (Currently amended) The <u>pharmaceutical</u> composition according to claim 1, wherein the composition is for intraperitoneal, subcutaneous, intranasal, intramuscular, intrathecal, sublingual, rectal, intravenous infusion, transdermal delivery or oral administration.
- 29. (Currently amended) The <u>pharmaceutical</u> composition according to claim 1, wherein the amount of Stofisopam, prodrug, or a pharmaceutically acceptable salt thereof is from <del>approximately</del> 10 mg to 1200 mg.
- 30. (Currently amended) The <a href="pharmaceutical">pharmaceutical</a> composition according to claim 1, wherein the amount of Stofisopam, prodrug, or a pharmaceutically acceptable salt thereof is from 50 mg to 600 mg.

- 31. (Currently amended) The <u>pharmaceutical</u> composition according to claim 1, wherein the amount of Stofisopam, prodrug, or a pharmaceutically acceptable salt thereof is from 100 mg to 400 mg.
- 32. (Currently amended) A method of administering a pharmaceutical composition according to claim 1, wherein the amount of comprising preparing the pharmaceutical composition comprising S-tofisopam, pro-drug or pharmaceutically acceptable salt administered is thereof and a pharmaceutically effective carrier and administering the pharmaceutical composition at a dose of less than 30 mg/kg.